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APPLICATION N	O	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/899,922 07/09/2001		07/09/2001	Amanda Johanne Kiliaan	BO 44633	5229
466	7590	06/13/2006		EXAMINER	
YOUNG	& THOM	PSON	DAVIS, RUTH A		
745 SOUT 2ND FLO	TH 23RD S' OR	TREET	ART UNIT	PAPER NUMBER	
	TON, VA	22202	1651		
			DATE MAILED: 06/13/2006		

Please find below and/or attached an Office communication concerning this application or proceeding.

	· · · · · · · · · · · · · · · · · · ·	Applicat	on No.	Applicant(s)	
		09/899,9	22	KILIAAN ET AL.	
Office Ac	Examin	r	Art Unit		
		Ruth A. D	avis	1651	
Th MAILING Period for Reply	DATE of this communic	cation appears on th	e cover sheet with the	correspondence ad	ddress
A SHORTENED STA WHICHEVER IS LON - Extensions of time may be after SIX (6) MONTHS from - If NO period for reply is spe - Failure to reply within the se Any reply received by the Co	TUTORY PERIOD FO NGER, FROM THE MA available under the provisions of the mailing date of this commu- cified above, the maximum state of or extended period for reply w office later than three months aft lent. See 37 CFR 1.704(b).	AILING DATE OF TO f 37 CFR 1.136(a). In no ex inication. utory period will apply and v iil, by statute, cause the app	HIS COMMUNICATIO rent, however, may a reply be ti rill expire SIX (6) MONTHS fror plication to become ABANDON	N. mely filed n the mailing date of this of ED (35 U.S.C. § 133).	
Status					
2a)⊠ This action is F 3)□ Since this appl	communication(s) filed [INAL. 2] cation is in condition for dance with the practice.	b)☐ This action is of allowance excep	t for formal matters, p		e merits is
Disposition of Claims					
4a) Of the abov 5) ☐ Claim(s) 6) ☑ Claim(s) <u>42-62</u> 7) ☐ Claim(s)		e withdrawn from co			
	n is objected to by the	Evaminar			
10) The drawing(s) Applicant may not Replacement drawing	n is objected to by the filed on is/are: ot request that any objection sheet(s) including claration is objected to	a) accepted or b tion to the drawing(s) the correction is requi	be held in abeyance. Se red if the drawing(s) is o	ee 37 CFR 1.85(a). ojected to. See 37 C	
Priority under 35 U.S.C	§ 119				
a) All b) So 1. Certified 2. Certified 3. Copies of applications.	nt is made of a claim forme * c) None of: copies of the priority of the certified copies of on from the Internation d detailed Office action	locuments have be locuments have be if the priority docum nal Bureau (PCT Ru	en received. en received in Applica ents have been receiv le 17.2(a)).	tion No red in this Nationa	l Stage
Attachment(s) 1) Notice of References Cit	ed (PTO-892)		4) Interview Summar		
2) D Notice of Draftsperson's	Patent Drawing Review (P1 tatement(s) (PTO-1449 or F		Paper No(s)/Mail I 5) Notice of Informal 6) Other:	Date	O-152)

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DETAILED ACTION

Applicant's amendment and response filed on April 3, 2006 have been received and entered into the case. Claims 61 - 62 are added; claims 42 - 62 are pending and have been considered on the merits. All arguments have been fully considered.

Claim Rejections - 35 USC § 103

- 1. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 2. This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).
- 3. Claims 42 48 and 51 62 are rejected under 35 U.S.C. 103(a) as being unpatentable over Horrobin, Fugh-Berman, Maggioni and Growdon.

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Applicant claims a method for treating a person having a vascular disorder and has or is at risk of developing unipolar depression, the method comprising orally administering a composition comprising (a) long chain polyunsaturated fatty acids (PUFAs) comprising omega-3 and omega-6 fatty acids in an amount of at least 350 mg/day, (b) phosphatidylethanolamine and phosphatidylcholine, and one of phosphatidylserine or phosphatidylinositol; in a ratio of 0.5 – 20; and (c) at least one compound selected from folate, vitamin B12, B6, magnesium, zinc. The composition further comprises hypericin or Withania somniera; 0.5 - 30g citrate; tryptophan or protein containing tryptophan; one of SAMe choline, betaine or copper; one of vitamin C, E, lipoic acid, selenium salt or carotenoids; ginkgo biloba extract; or vitamin D. The omega-3 fatty acids are selected from eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) and the omega-6 fatty acids are selected from arachidonic acid (ARA) and dihomogamma linolenic acid (DGLA). The (c) portion contains both folate and vitamin B6. Specifically, the composition comprises at least 120 mg long chain PUFAs, 200mg phospholipids, 200 micrograms folate, 0.1 mg hypericin or 100 mg W. somnifera, and 500 mg citrate. The phospholipids are in the amount of 1 g/day. Applicant additionally claims a method for treating a person having a vascular disorder and has or is at risk of developing unipolar depression, the method comprising administering a composition comprising (a) 350 mg of long chain PUFAs wherein the omega-3 fatty acids are EPA and DHA, and the omega-6 fatty acids are AA and DHGLA at a ratio of 2.5 - 5.5:1; (b) at least 2 phospholipids selected from phosphatidylethanolamine and phosphatidylcholine, and one of phosphatidylserine or phosphatidylinositol; (c) a compounds selected from folate, vitamin B12, B6, magnesium, zinc. Specifically, at least 20 mg EPA, 50 mg DHA, 50 mg AA, 200 mg phospholipids, 200 mg folate, 0.2 mg hypericin or 500 mg

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W.somnifera, 100 mg Mg, 5 mg Zn, 2 mg vitamin B6, 2 micrograms B12 and 1 g citrate.

Applicant finally claims the method wherein the composition comprises 350 mg long chain PUFAs; at least 2 phospholipids selected from phosphatidylethanolamine, phosphatidylcholine, phosphatidylserine or phosphatidylinositol; a compound selected from folate, vitamin B12, B6, magnesium, zinc; and 4 – 40 micrograms of vitamin D3.

Horrobin teaches compositions and methods for treating depression and anxiety, the compositions comprising DHA (p.3,4), ascorbic acid (vitamin C), vitamin E, beta carotene, selenium, zinc, and vitamin B6 (p.5,7, claims). The composition may further comprise EPA, DGLA, and AA, (p.6). Specifically, dosages of at least 350 mg DHA are combined with 250 – 2000 mg of the other named fatty acids (p.6). Examples further include vitamins of the B group, vitamin D, folic acid (folate), magnesium, and lipoic acid (examples).

Fugh-Berman teaches St. John's Wort, or hypericine (p.713), ginkgo biloba (p.715-16), vitamin B12, folate (p.721), SAMe, and tryptophan (p.722) improve depression and symptoms thereof.

Growdon teaches methods for treating depression and related disorders comprising administering lecithin (consists of phosphatidylcholine, phosphatidylethanolamine, and phosphatidylinositol, see attached "Soy Lecithin Fact Sheet") (abstract, col. 1-3).

Maggioni teaches phosphatidylserine for treating depression and symptoms thereof (abstract).

The above references do not teach a composition comprising all of the combined ingredients. However, at the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to combine the instant ingredients for their known benefit, as disclosed

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by the cited references above, since each is well known in the art for their claimed purpose. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated to combine each of the claimed ingredients together with a reasonable expectation for successfully treating depression. Since each of the claimed ingredients were known to have the same, claimed, therapeutic effects, one in the art would know that they are result effective variables. Thus, while references do not teach the specific amounts or ratios as claimed, it would have been well within the purview of one of ordinary skill in the art to optimize amounts and/or ratios as a matter of routine experimentation. In addition, although the references do not specifically teach inclusion of citrate, citrate was a well known stabilizer and synergist with various vitamins (as admitted by applicant, specification p.5). It would have been obvious to one of ordinary skill in the art to include citrate as a matter of routine practice at the time the claimed invention was made. This rejection is based on the well established proposition of patent law that no invention resides in combining old ingredients of known properties where the results obtained thereby are no more than the additive effect of the ingredients, In re Sussman, 1943 C.D. 518. It is noted that where general conditions of a claim are disclosed in the prior art, it is not inventive to disclover the optimiun or workable ranges by routine experimentation. The normal desire of one in the art to improve upon what is already generally known provides the motivation to determine where in a disclosed set of ranges is the optimum combination (MPEP 2144)

Thus, the invention as a whole is *prima facie* obvious over the references, especially in the absence of evidence to the contrary.

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4. Claim 49 is rejected under 35 U.S.C. 103(a) as being unpatentable over Horrobin, Growdon and Pollack.

Applicant claims a method for treating a person having a vascular disorder and has or is at risk of developing unipolar depression, the methodcomprising orally administering a composition comprising (a) long chain polyunsaturated fatty acids (PUFAs) comprising omega-3 and omega-6 fatty acids in an amount of at least 350 mg/day, (b) phosphatidylethanolamine and phosphatidylcholine, and one of phosphatidylserine or phosphatidylinositol; in a ratio of 0.5 – 20; and (c) at least one compound selected from folate, vitamin B12, B6, magnesium, zinc. The composition further comprises copper, with a ratio of zinc to copper of 5 – 12:1.

Horrobin teaches compositions and methods for treating depression and anxiety, the compositions comprising DHA (p.3,4), zinc, and vitamin B6 (p.5,7, claims). The composition may further comprise EPA, DGLA, and AA, (p.6). Specifically, dosages of at least 350 mg DHA are combined with 250 – 2000 mg of the other named fatty acids (p.6). Examples further include vitamins of the B group, folic acid (folate), magnesium, and lipoic acid (examples).

Growdon teaches methods for treating depression and related disorders comprising administering lecithin (consists of phosphatidylcholine, phosphatidylethanolamine, and phosphatidylinositol, see attached "Soy Lecithin Fact Sheet") (abstract, col. 1-3).

Pollack teaches methods for treating depression comprising administering compositions comprising vitamin B6 (pyridoxine), copper and magnesium (abstract, claims 8-14).

The above references do not teach a composition comprising all of the combined ingredients. However, at the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to combine the instant ingredients for their known benefit, as disclosed

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by the cited references above, since each is well known in the art for their claimed purpose. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated to combine each of the claimed ingredients together with a reasonable expectation for successfully treating depression. Since each of the claimed ingredients were known to have the same, claimed, therapeutic effects, one in the art would know that they are result effective variables. Thus, while references do not teach the specific amounts or ratios as claimed, it would have been well within the purview of one of ordinary skill in the art to optimize amounts and/or ratios as a matter of routine experimentation. This rejection is based on the well established proposition of patent law that no invention resides in combining old ingredients of known properties where the results obtained thereby are no more than the additive effect of the ingredients, In re Sussman, 1943 C.D. 518. It is noted that where general conditions of a claim are disclosed in the prior art, it is not inventive to disclover the optimiun or workable ranges by routine experimentation. The normal desire of one in the art to improve upon what is already generally known provides the motivation to determine where in a disclosed set of ranges is the optimum combination (MPEP 2144)

Thus, the invention as a whole is *prima facie* obvious over the references, especially in the absence of evidence to the contrary.

5. Claim 50 is rejected under 35 U.S.C. 103(a) as being unpatentable over Horrobin, Growdon and Takeda.

Applicant claims a method for treating a person having a vascular disorder and has or is at risk of developing unipolar depression, the method comprising orally administering a

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composition comprising (a) long chain polyunsaturated fatty acids (PUFAs) comprising omega-3 and omega-6 fatty acids in an amount of at least 350 mg/day, (b) phosphatidylethanolamine and phosphatidylcholine, and one of phosphatidylserine or phosphatidylinositol; in a ratio of 0.5 – 20; and (c) at least one compound selected from folate, vitamin B12, B6, magnesium, zinc. The composition further comprises at least one of carnitine, B1, B5 or CoEnzyme Q10.

Horrobin teaches compositions and methods for treating depression and anxiety, the compositions comprising DHA (p.3,4), zinc, and vitamin B6 (p.5,7, claims). The composition may further comprise EPA, DGLA, and AA, (p.6). Specifically, dosages of at least 350 mg DHA are combined with 250 – 2000 mg of the other named fatty acids (p.6). Examples further include vitamins of the B group, folic acid (folate), magnesium, and lipoic acid (examples).

Growdon teaches methods for treating depression and related disorders comprising administering lecithin (consists of phosphatidylcholine, phosphatidylethanolamine, and phosphatidylinositol, see attached "Soy Lecithin Fact Sheet") (abstract, col. 1-3).

Takeda teaches compositions for treating depression comprising carnitine and vitamin B1 (abstract).

The above references do not teach a composition comprising all of the combined ingredients. However, at the time of the claimed invention, it would have been obvious to one of ordinary skill in the art to combine the instant ingredients for their known benefit, as disclosed by the cited references above, since each is well known in the art for their claimed purpose. Moreover, at the time of the claimed invention, one of ordinary skill in the art would have been motivated to combine each of the claimed ingredients together with a reasonable expectation for successfully treating depression. Since each of the claimed ingredients were known to have the

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same, claimed, therapeutic effects, one in the art would know that they are result effective variables. Thus, while references do not teach the specific amounts or ratios as claimed, it would have been well within the purview of one of ordinary skill in the art to optimize amounts and/or ratios as a matter of routine experimentation. This rejection is based on the well established proposition of patent law that no invention resides in combining old ingredients of known properties where the results obtained thereby are no more than the additive effect of the ingredients, *In re Sussman*, 1943 C.D. 518. It is noted that where general conditions of a claim are disclosed in the prior art, it is not inventive to disclover the optimiun or workable ranges by routine experimentation. The normal desire of one in the art to improve upon what is already generally known provides the motivation to determine where in a disclosed set of ranges is the optimum combination (MPEP 2144)

Thus, the invention as a whole is *prima facie* obvious over the references, especially in the absence of evidence to the contrary.

Response to Arguments

Applicant first argues that the references do not teach the claimed amounts and ratios of phospholipids; that there is no motivation to combine and optimize the amounts of ingredients; that the ingredients are not result effective variables; that it is unexpected that the claimed diet is better than a diet without the supplement for treating depression, as evidenced by the declaration submitted on December 10, 2004; and that supporting references do not remedy the deficiencies

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Regarding applicant's assertion that it would not be obvious to optimize the amounts and ratios of the instant ingredients, it is reiterated that it would have been well within the purview of one of ordinary skill in the art to optimize the amounts and ratios of ingredients since they were known to have the claimed activity. It is specifically noted that Horrobin teaches a wide range of effective amounts of phospholipids (p.4,6,7) indicating that one in the art would recognize the ingredients as result effective variables. Thus one in the art would know to optimize the amount of such active ingredients. It is further noted that the ratios do not appear to impart unexpected advantages or results to the claimed composition.

Regarding applicant's claim that the instant composition is unexpectedly better, it is reiterated that the compositions described in the affidavit are not the same as the claimed composition, thus is not commensurate in scope with the claimed method. In order to provide convincing evidence of an unexpected advantage or benefit, the evidence must be commensurate in scope with the claimed composition and method.

Conclusion

6. Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, THIS ACTION IS MADE FINAL. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE

MONTHS from the mailing date of this action. In the event a first reply is filed within TWO

MONTHS of the mailing date of this final action and the advisory action is not mailed until after

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the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Ruth A. Davis whose telephone number is 571-272-0915. The examiner can normally be reached on M-F 7:00 - 2:30pm.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Michael Wityshyn can be reached on 571-272-0926. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

June 9, 2006 AU 1651 RUTH A. DAVIS
PATENT EXAMINER